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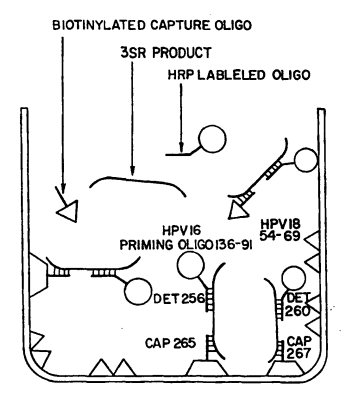
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(54) Title: HUMAN PAPILLOMAVIRUS DETECTION ASSAY

(57) Abstract

A two-step nucleic acid hybridization probe assay for certain types of human papilloma virus (HPV) associated with cervical cell dysplasia and malignancy comprises a fluid phase capture hybridization step in which amplified specific gene E6/E7 messenger RNA from a biological specimen is hybridized to a biotinylated capture reagent to form a complex, attachment of the capture reagent complex to a solid phase by reaction with immobilized streptavidin, a second hybridization step in which a virus type-specific enzyme-conjugated detection probe hybridizes with the complexed amplified messenger RNA, and detection of the complexed detection probe by color or fluorophor production following a wash of the solid phase and addition of an appropriate chromogenic or fluorogenic substrate. The assay has enhanced sensitivity compared to conventional tests and is specific for actual expression of HPV oncogenes in cervical specimens, and not merely indicative of viral presence.



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HUMAN PAPILLOMAVIRUS DETECTION ASSAY

BACKGROUND OF THE INVENTION

Human papillomaviruses (HPVs) are a heterogeneous group 5 of DNA viruses associated with a variety of proliferative lesions of the epithelium. Many of these lesions are benign such as those associated with HPV 6 and HPV 11, and are considered causative of such conditions as warts, and condylomas (see Gissman, Canc. Surv., 3: 161 (1984)). However, epidemiological and molecular studies implicate several high risk types that infect the genital tract associated with dysplasia and sometimes progress to cervical cancer (see, for example, Durst. et al., PNAS, 80: 3812 (1983)). High risk HPV types are predominately HPV 16 and HPV 18, with HPV 31, HPV 33, and HPV 35 being of lesser significance. More recently, another HPV type associated with malignancy, HPV 44, has been identified (Lorincz, U.S. Patent No. 4,849,331).

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HPV of any type is generally found in extremely low numbers in biological specimens. Therefore, molecular techniques must be performed for amplifying nucleic acid viral markers from very low copy number in a specimen to detectable levels. Polymerase chain reaction (PCR) has been utilized to amplify HPV viral DNA in this manner, as disclosed in WO 90/02821, and Shibata, et al., J. Exp. Med., 167: 225 (DATE). Other applications of PCR to HPV diagnostics are Maitland, et al., May WO 94/26934

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1988. Seventh International Papillomavirus Workshop, Abstract, p. 5 and Campione-Piccardo, et al., May 1988, Seventh International Papillomavirus Workshop. One major problem with PCR amplification of HPV is that these viruses are detectable as fortuitous passengers in a significant percentage of healthy women showing no evidence of any benign of malignant pathology. Percentage estimates of such passenger presence range 10% (see U.S. Patent No. 4,983,728) to as high as 60%. Detection of HPV per se is thus of limited diagnostic value.

Many nucleic acid-based assays utilize the well-known sandwich configuration in a heterogeneous format. In this format a capture oligonucleotide is chemically conjugated to a solid support such as a microtiter well or bead, the sample is added, and the target nucleic acid having base homology to capture oligonucleotide is allowed to hybridize. After a wash (phase separation), a detection oligonucleotide hybridizes, and after a second wash to remove unhybridized detection oligonucleotide, the amount of tracer or reporter is measured, or the signal generating means produces a signal. For the details of such assays, refer to Ranki, U.S. Patent No. 4,486,539 and U.S. Patent No. 4,731,325. The basic problem with such sandwich assays is relatively low capture efficiency on the solid support, which may profoundly reduce sensitivity of the assay.

SUMMARY OF THE INVENTION

It is an object of this invention to provide a specific assay for HPV infections associated with cervical dysplasia and cellular transformation to malignancy. In achieving this object, it is essential to first amplify to detectable levels only the messenger RNA (mRNA) expressed from oncogene regions (genes E6/E7) of HPV types implicated in malignant or pre-malignant cervical lesions. This not only restricts detection to malignant and pre-malignant HPV types, but also distinguishes actual oncogene expression from mere passenger presence of virus.

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It is a further object to provide a highly sensitive assay for HPV having a high capture efficiency in the initial capture hybridization step. This is important because in situations in which the patient specimen contains very low copy number of viral mRNA, amplification may not occur to a level high enough for detection unless the assay itself is sensitive.

It is a still further aspect of the invention to provide reagents such as primer families for optimally efficient amplification, and probes which anneal to their targets under stringent conditions to give high selectivity and specificity. Finally, the invention contemplates a kit comprising these reagents, buffers, sample preparation solutions, solid supports, and reaction vessels.

In accordance with the assay of the present invention, a patient specimen suspected of containing messenger RNA encoded by at least one type of HPV associated with cervical dysplasia, malignant cells, or pre-malignant cells is

(1) subjected to nucleic acid amplification by self sustained sequence replication utilizing two primers separated by at least ten nucleotides, at least one such primer containing a transcriptional promoter,

annealing the first such primer to its complementary sequence on the target region messenger RNA, extending the 3' end of the primer by action of a strand-extending polymerase in the presence of cofactors and nucleotide triphosphates,

digesting the RNA strand of the nascent RNA/DNA duplex with an enzyme having exogenous or endogenous RNAse H activity,

annealing the second such primer to its complementary sequence on the resultant single stranded cDNA, primer extending the 3' end of the primer by action of a strand-extending polymerase,

transcribing the double stranded DNA with a transcriptase in the presence of nucleoside triphosphates, and

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repeating the amplification utilizing the newly synthesized transcripts as new targets,

- (2) hybridizing in solution amplified messenger RNA to a free biotinylated reagent capture probe having a sequence complementary to a first segment of the amplified RNA to form a reagent capture complex,
- (3) attachment of the capture complex to a solid phase by reaction of the biotin residue of the capture probe with streptavidin bound to the surface of the solid phase,
- (4) washing the bound complex to remove unbound and10 unreacted reagents,
 - (5) hybridizing a virus type-specific enzyme-conjugated detection probe having a sequence complementary to a second segment of the amplified RNA not overlapping the sequence of the first such RNA segment to form a solid phase-bound capture probe-target sequence-detection probe complex,
 - (6) washing the complex to remove unhybridized detection probe, and
 - (7) adding a fluorogenic or chromogenic enzyme substrate and reacting the conjugated enzyme to produce a detectable fluorophor or chromogen.

The present invention is also directed to certain primer families and selected probes for use in the HPV detection assay, and to kits for conveniently providing reagents to users.

25 BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1: HPV 16 genome organization. Transcription proceeds clockwise from the P_{97} promotor. A_E and A_L are the polyadenylation sites for the early and late transcripts.

Figure 2: Sequence of HPV 16. The primers are indicated by underlines. Boxes indicate splice donor and acceptor sequences.

Figure 3: Sequence of HPV 18. Sequences of HPV 18 primers are indicated by underlines. Boxes indicate splice donors and acceptor sequences.

Figure 4: HPV 16 primer families. A variety of primers were tested by the ability to amplify total RNA from SiHa cells (infected with HPV 16). The reactions contained 10% DMSO and 15% sorbitol. The primers are indicated on the autoradiogram.

Figure 5: The effect of increasing the RNAse H concentration using HPV 16 primer families.

10 Figure 6: HPV 16 primer sensitivity. Total RNA is titrated from 1, 0.1, 0.01, 0.001 attomoles of specific E6-7 RNA isolated from SiHa RNA. p. 32. N5.

Figure 7: Primer sensitivity using cells which contain HPV 18 DNA. From right to left is 104 to 10 cells. p34 N4.

Figure 8: An autoradiogram slotting 3SR reaction products.

A RNAse titration was performed using primers 32-54 which amplified HPV 18 RNA.

Figure 9: Autoradiogram of a 3SR reaction using primers 32-54 containing different additives. The additives (left to right) were 10% DMSO, 10% polyethylene glycol and 10% glycerol. The cross reactivity using primers 29-15 using SiHa cell using these additives were included to determine if there was any cross reactivities of the reactions.

Figure 10: Autoradiogram of a 3SR reaction comparing primers 32-54 and 69-54. The 3SR reaction using primers 69-54 contained either no additives (column 1) or 15% sorbitol (column 2). The reactions using Primers 32-54 contained 10% polyethylene glycol (column 3). From top to bottom was a titration of RNAse H, 1-3 units per reaction.

Figure 11: Co-amplification. Lane A used primers 136-73 (HPV 16), Lane B used primers 136-91 (HPV 16) amplifying 5 amol of SiHa RNA using decreasing amounts of DMSO/sorbitol mixture. Lane C from top to bottom: 136-73 (HPV 16) and 54-69 (HPV 18), 136-91 and 54-69, and 54-69 amplifying a mixture of 5 amol of SiHa cell (infected with HPV 16)

and HeLa cell (infected with HPV 18) RNA. Duplicate blots were prepared and probed with an HPV 18 specific probe (59) and an HPV 16 specific probe (98).

Figure 12: HPV 16 plate optimization. Capture 245 temperature optimum. Absorbance values using CAP245 at different temperature ranges: 30°C, 40°C, 50°C, 60°C and 70°C. Each line represents a different detectors; DET 251, DET 252, and DET 254.

Figure 13: HPV 16 plate optimization. Capture 250 temperature optimum. Absorbance values using CAP250 at different temperature ranges: 30°C, 40°C, 50°C, 60°C and 70°C. Each line represents a different detectors; DET 251, DET 252, and DET 254.

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Figure 14: Detector hybridization optimum using CAP 245. Detectors were hybridized using different temperature ranges: 30°C, 40°C, 50°C, 60°C and 70°C. Each line represents different detectors: DET 98, DET 251, DET 252, and DET 254.

Figure 15: Detector hybridization optimum using CAP 250. Detectors were hybridized using different temperature ranges: 30°C, 40°C, 50°C, 60°C and 70°C. Each line represents different detectors: DET 98, DET 251, DET 252, and DET 254.

Figure 16: HPV 16 plate assay. A comparison of captures 245, 250, and 253 using DET 98, DET 251, DET 252, and DET 254. Each capture was hybridized to the 3SR product at 50°C. The detectors were hybridized at room temperature.

Figure 17: HPV 16 detector performance. A comparison of all the detector oligos for HPV 16 using CAP 250. The detector names are listed in the bottom of each figure.

Figure 18: A comparison of detector lengths using CAP 250 in the enzyme probe assay. DET 256 is a 17mer oligo and DET 257 is a 15mer oligo. The sequence was identical except that 2 bases were omitted for DET 257.

Figure 19: A comparison in absorbance values using different additives in the capture buffer. From left to right are duplicate wells using

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DET 255, DET 98 and DET 256. Columns 1-6 are 3SR products using primers 96-91. Columns 7-12 are 3SR products using primer 137-91 using different detectors. The additives are indicated on the left of the absorbance values. Rows 1 and 2 are plus and minus templates using 5% polyethylene glycol. Rows 3 and 4 are plus and minus templates using 1% BSA. Rows 5 and 6 are plus and minus templates using 5% PEG, 1% BSA. Rows 7 and 8 are the standard hybridization buffer using 0.1% polyvinylpyrrolidone, 5X SSC.

Figure 20: A comparison in absorbance values using different additives in the detection buffer. From left to right using different detectors: DET 256, DET 98, and DET 255. Columns 1, 5, and 9 contained the standard hybridization buffer 30% glycerol, 0.1% PVP, 1% BSA and 5X SSC. Columns 2, 6, and 10 contained 5% PEG, 0.1% PVP, and 5X SSC as the hybridization buffer. Columns 3, 7, and 11 contained 1% BSA, 0.1% PVP and 5X SSC as the hybridization buffer. Columns 4, 8, and 12 contained 5% PEG, 1% BSA, 0.1% PVP, and 5X SSC as the hybridization buffer. Rows A and B are plus and minus templates using primers 96-91 which amplify SiHa RNA. Rows C and D is plus and minus template using primers 136-91 which amplify SiHa RNA.

Figure 21: Different primers sets which amplify HeLa RNA (HPV 18). Primers are noted on the autoradiogram.

Figure 22: Comparison of capture oligos for HPV 18 using the enzyme probe assay. The 3SR product was amplified from HeLa RNA using primer 54-69. Column 1 is substrate only. Columns 2 and 3 are plus and minus templates using capture 56. Columns 4 and 5 is plus and minus templates using capture 267. Rows indicate different detectors. Row A DET 59, Row B DET 260, Row C DET 262, Row D DET 268, Row E DET 269, and Row F DET 270.

Figure 23: Comparison of capture oligos for HPV 16 and HPV 30 18 using the enzyme probe assay. The 3SR product was a co-amplification from HeLa and SiHa RNA using primers 136-91 (HPV 16) and 54-69 (HPV 18).

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Figure 24: HPV 16 and HPV 18 EPA. The absorbance levels of a typical specimen. HPV 16 and HPV 18 were co-amplified using primers 136-91 and 54-69. CAP 265 and CAP 267 were added and allowed to hybridize. The reaction was added to two microwells and detected using a type specific oligo DET 256 and HPV 16 and DET 260 for HPV 18.

Figure 25: Schematic of the Enzyme Probe Assay. The capture oligo hybridizes to the amplified 3SR product either HPV 16 or HPV 18. The complex is detected using HRP labeled oligonucleotide.

Figures 26 and 27: Autoradiographs of amplification products comparing yields of reaction performed at 50°C and at 42°C.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

Figure 1 is a schematic drawing showing a generalized HPV 16 genome. The heavy concentric lines indicate open reading frames. Figures 2 and 3 locate the splice donor and acceptors for HPV 16 and 18 genes (indicated by boxes around the terminal two bases involved in the splice in the E6/E7 region). The portion of the HPV 16 and 18 viral genomes coding for E6/E7 polypeptides are identified in the Sequence Listing as SEQ. ID. Nos. 1 and 2 respectively. This is a significant region of the genome since the proteins encoded are thought to be involved in degradation of the p53 suppressor protein, which regulates cell growth. Loss of p53 function is associated with malignancy. Thus, expression of E6/E7 is diagnostic for cervical cancer or pre-malignant states.

In the expression of the E6/E7 region, splicing at the positions indicated in the figures occurs at substantial but unknown frequency. In designing primers for amplification of mRNA targets transcribed from this region, it is therefore important to make certain that all primer pairs lie outside the portion of the transcript from which the splice leads to excision of an mRNA fragment. Typical primers selected are illustrated in figures 2 and 3.

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Since the rationale of the assay of the present invention is to detect only gene products produced in cells actually expressing genes

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E6/E7, self-sustained sequence replication (3SR) is the amplification method of choice. Polymerase chain reaction amplifies DNA, and while it may detect the presence of virus with great sensitivity, it is unsuitable for detecting gene expression. The method of 3SR is fully described in Gingeras, et al., Ann. Biol. Clin., 48: 498 (1990), Guatelli, et al., PNAS, 87: 1874 (1990), and WO 90/06995. The methods described therein are followed herein except as noted, and define the procedure to be followed in the practice of the present invention. The general 3SR amplification procedure as set forth in Gingeras et al. and Guatelli et al. involves the following steps: One hundred-microliter 3SR amplification reactions contained the target RNA, 40 mM Tris-HCl at pH 8.1, 20 mM MgCl₂, 25 mM NaCl, 2 mM spermidine hydrochloride, 5 mM dithiothreitol, 80 μg/ml bovine serum albumin, 1 mM dATP, 1 mM dCTP, 1 mM dGTP, 1 mM dTTP, 4 mM ATP, 4 mM CTP, 4 mM GTP, 4 mM UTP, and 250 ng of each selected oligonucleotide primer. After heating at 65°C for 1 minute and cooling at 37°C for 2 minutes, 30 units of AMV reverse transcriptase, 100 units of T7 RNA polymerase, and 4 units of E. coli RNase H were added to each reaction. All reactions were incubated at 37°C for 1 hour and stopped by placing the reaction on ice.

In general, 3SR is carried out as follows on HPV specimens: samples are obtained by vaginal lavage or cervical scrape. Messenger RNA is released by treatment with chaotrophic/phenol reagents and precipitated conventionally with ethanol. A preferred one step extraction utilizes RNAzol B (Cinna/Tiotecx Laboratories, Inc.) according to the manufacturer's instructions. The RNA is then dissolved in 3SR buffer, together with nucleotide and nucleoside triphosphates, primers, enzymes, and cofactors to carry out 3SR amplification. Reagents were obtained as follows:

30 Primer Oligonucleotides

All oligonucleotides may be synthesized on a commercially available synthesizer such as a Milligen 8700 DNA synthesizer.

Oligonucleotides which contained a 5' biotin may be synthesized using a biotin phosphoramidite (Glenn Research). Oligonucleotides which contain a 3' biotin may be synthesized using control pore glass containing a protected biotin (Glenn Research). Oligonucleotides which contain a 3' amine are conveniently synthesized using a amino-on control pore glass column (Glenn Research). Below is a list of oligonucleotides used in the development of HPV 16/18 enzyme probe assay of the present invention. All of the sequences are from left to right 5' to 3'. The oligonucleotide primers are also listed in the Sequence Listing as SEQ. ID. Nos. 3-31.

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	SEQ. ID. No.	Primer Pro	<u>bes</u>
	3	HPV15:	AAT TTA ATA CGA CTC ACT ATA GGG
			AGC TIT TCT TCA GGA CAC AGT GGC
			T
15	4	HPV19:	AAT GTT TCA GGA CCC ACA GGA GC
	5	HPV20:	GAA TGT GTG TAC TGC AAG CAA
			CAG
	6	HPV29:	ATG CAC AGA GCT GCA AAC AAC TA
	7	HPV32:	CAC TTC ACT GCA AGA CAT AGA A
20	8	HPV48:	AAT TTA ATA CGA CTC ACT ATA GGG
			ATG TGT CTC CAT ACA CAG AGT C
	9	HPV53:	GAA TGT GTG TAC TGCC AAG CAA
			CAG
	10	HPV54:	AAT TTA ATA CGA CTC ACT ATA GGG
25			AAA GGT GTC TAA GTT TTT CTG CTG
			G
	11	HPV69:	CTG AAC ACT TCA CTG CAA GAC
	· 12	HPV73:	CAG TTA TGC ACA GAG CTG CAA AC
	13	HPV74:	GTT ATG CAC AGA GCT GCA AAC AA
30	14	HPV77 :	CAA GCA ACA GTT ACT GCG AC
	15 .	HPV89:	AGC AAC AGT TAC TGC GAC GT
	16	HPV90:	GCA CAG AGC TGC AAA CAA CTA TA

	17	HPV91:	ACA GAG CTG CAA ACA ACT ATA CA
	18	HPV92:	AAT TTA ATA CGA CTC ACT ATA GGG
			ACT TTT CTT CAG GAC ACA GTG GCT
			TTT
5	19	HPV93:	AAT TTA ATA CGA CTC ACT ATA GGG
			ATT TGC TTT TCT TCA GGA CAC AGT
			GG
	20	HPV94:	AAT TTA ATA CGA CTC ACT ATA GGG
			ATC TTT GCT TTT CTT CAG GAC ACA
10			GT
	21	HPV95:	AAT TTA ATA CGA CTC ACT ATA GGG
			ATG TCT TTG CTT TTC TTC AGG ACA
			CA
	22	HPV96:	AAT TTA ATA CGA CTC ACT ATA GGG
15			AGA TGT CTT TGC TTT TCT TCA GGA
			CA
	23	HPV101:	AGA GCT GCA AAC AAC TAT ACA TG
	24	HPV106:	AAT TTA ATA CGA CTC ACT ATA GGG
			ATT CAT GCA ATG TAG GTG TAT CTC
20			С
	25	HPV107:	AAT TTA ATA CGA CTC ACT ATA GGG
			ATA TTC ATG CAA TGT AGG TGT ATC
			T
	26	HPV118:	AGC TGC AAA CAA CTA TAC ATG AT
25	27	HPV120:	AAT TTA ATA CGA CTC ACT ATA GGG
	•		ATG CAA TGT AGG TGT ATC TCC ATG
			С
	28	HPV129:	AAT TTA ATA CGA CTC ACT ATA GGG
			AAA TGT AGG TGT ATC TCC ATG CAG
30	29	HPV131:	AAA CAA CTA TAC ATG ATA TAA TA

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	30	HPV136:	AAT TTA ATA CGA CTC ACT ATA GGG
			AAT GTA GGT GTA TCT CCA TGC ATG
			A
	31	HPV137:	AAT TTA ATA CGA CTC ACT ATA GGG
5			ATG TAG GTG TAT CTC CAT GCA TGA
			Т

Primer selection for high level amplification is basically a directed trial and error process. To define a first set of primers a span of 400 bases (with beginning and ending sites outside the spliced region) was selected by designating the first 10-30 nucleotides at the 5' end of the E6 gene beginning with the ATG codon and counting off 400 bases, then selecting as primers the next 10-30 bases. Note that for each pair, at least one of the primers must contain a promoter for transcription. The bacteriophage T7 RNA polymerase binding site (SEQ. ID. No. 44), AAT TTA ATA CGA CTC ACT ATA GGG A, is preferred because of its strength and specificity.

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The primer pairs are tested for their amplification efficiency. To optimize, the second primer position is held stationary and the first primer is moved arbitrarily 20 bases towards the second (thereby decreasing the interprimer span, e.g. the bases between the position of the 3' end of the first primer and the 5' end of the second primer, by 20 bases to 380 bases). Fine tuning is accomplished by walking the primers from the best pairings by 2-5 base jumps.

Primer families. Figure 4 gives primer families that amplify the HPV 16 E6-7. All primers amplified total RNA isolated from the SiHa cell line which contain the HPV 16 transcripts. The reaction conditions include 7mM rNTPs, 1mM dNTPs, 40mM Tris pH 8.1, 30mM MgCl₂ 20mM KCl, 50mM dithiothreitol, 20 mM spermidine, 10% DMSO, 15% sorbitol, and 15pmol each priming oligonucleotide. After pre-warming each tube at 42°C for 5 minutes 30 units of AMV-RT, 2 units RNAse H, and 250 units of T7 RNA polymerase were added as a cocktail to each reaction. The

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reaction was allowed to proceed for one hour at 42°C. A sample of the 3SR reaction was slotted onto nitrocellulose. The nitrocellulose was baked for 45 minutes and then hybridized for 45 minutes using a type specific detection oligo. An autoradiogram was generated by exposing the nitrocellulose to film for 45 minutes at -70°C. The primer family for 120 is 29 and 90. The primer family for 15 is 19, 20, 77, 53, and 89. The primer family for primer 129 is 29, 74, 73, 118, 130, and 131. The primer family for primer 136 is 91, 29, 90, 74, 73, 130, 131, and 118. The primer family for primer 137 is 29, 90, 74, 73, 131, and 118.

Figure 5 illustrates the effect of titrating the RNAse H HPV 16 primer families. The 3SR reaction conditions are identical as described in figure 4 except the DMSO and sorbitol were omitted from the reaction. Ten microliters were slotted onto nitrocellulose then baked and probed with a type specific detection oligo (HPV55). The primer family for primer 93 is 73 and 91. The optimal RNAse H needed for the reaction using these two primer pairs is between 1 and 2 units. The primer family 95 is 101 and 91. These primer sets do not appear to be sensitive to different RNAse H concentrations. A single primer set was defined for primer 92; 92-91, primer 94; 94-91, and primer 85; 85-77. The primer family for primer 96 is 73 and 91. All of these primer sets amplify optimally using between 2 and 3 units of RNAse H. The sensitivity of primers 96-73, 96-91, and 94-91 were tested using a titration of E6-7 isolated from SiHa cells. Once each primer set has been defined and optimized the sensitivity can be measured by amplifying decreasing amounts of RNA from control cells (figure 6). The 3SR reaction conditions are identical to those described in figure 4 except, using primers 96-73 the DMSO was included and the sorbitol was omitted, and using primers 94-91 only 10% sorbitol was included.

Figures 7-10 describe the primers used to amplify HPV 18 E6-7. The primer family for primer 54 is 32, 69, and 70. Primers 48 and 32 also amplify HeLa RNA. Primers 54-32 and 54-48 both require the addition of additives 10% polyethylene glycol or DMSO and sorbitol to the 3SR reaction. Primers 54-69 do not require the addition of additives for

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successful amplification. Additional primer families for primer 214 is 69, 244, 214, and 70 all which require additives to the amplification reaction.

Co-amplification. Once primers have been selected for both HPV 16 and HPV 18 a co-amplification of both targets is required for clinical use. Co-amplification is required because only a single specimen is obtained. This can be done not only for HPV 16 or HPV 18, but also can be applied to a plurality of HPV types including but not limited to HPV 31, 33, and 35, as well as any other types that prove to be oncogenic. It is not practical to split a single specimen for two independent reactions. Figure 11 is a duplicate blot which is probed with a 16 and 18 type specific detection probe. Lane C demonstrates the cross reactivity of amplifying two independent targets.

Capture and Detection Probes. Because it is impractical to incubate the plate in elevated temperatures the detector should produce maximum signal at room temperature. Many times uneven temperatures across a microwell can cause differences in hybridization thereby causing variability of absorbance values. The format of the plate affects the performance of the assay. Incubating both capture and detector probes simultaneously rather than capturing the 3SR product first and detecting in a separate incubation step affects the relative OD values. There are disadvantages of co-incubation of both capture and detection probes. In high template concentration, the 3SR reaction produces very high product concentrations. When the capture is incubated to the target in one step then applied to the microwell and allowed to bind, excess target is subsequently washed away. The detection probe is then applied which only hybridizes to the capture 3SR target.

When designing capture oligonucleotide sequences, defining the hybridization temperatures is critical to the performance of the assay. Figures 12 and 13 define the optimum temperature of hybridization for HPV 16 capture oligonucleotide. The 3SR product is diluted 1:10,000 to reduce the absorbance levels thereby allowing differences of different detection probes to become more pronounced. The hybridization reaction

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contain 50 µl of the diluted 3SR product in 0.1% PVP, 2X SSC, and 4 pmol capture oligonucleotide. The reaction was incubated at different temperatures ranging from room temperature to 70°C. The reaction proceeded in the microwell for 20 minutes and the well washed 3 times with 2X SSC (0.6 M NaCl, 0.06 M Na citrate pH 7.0), 0.05% Tween 20®, and 0.01% Thimersol™. The detection probe was added and incubated for 30 minutes at room temperature. The microwell was again washed 3 times with 2X SSC, 0.5% Tween 20, and 0.01% Thimersol. Substrate for the horseradish peroxidase enzyme, 3′, 3′, 5′, 5′, tetra methyl benzidine and hydrogen peroxide was added to each well and allowed to develop for 15 minutes at room temperature. The reaction was stopped by the addition of 1 M phosphoric acid and read at 450 nm.

The optimum temperature of hybridization for capture 245 is between 50°C and 60°C. The signal remains relatively constant at 70°C but thermal degradation of the RNA is a concern at this temperature. Capture 250 hybridization optimum is between 50°C and 60°C. A variety of detection probes should be tested because the optimum temperatures for hybridization of the detection probes must be empirically determined. Once the capture oligo temperature optimum has been defined, the same experiments must be repeated using different probes.

Best Mode. Figures 14 and 15 define the detector optimum. CAP 250 and CAP 245 produced the highest absorbance values when hybridizing DET 251 at room temperature. The reaction was performed as described in figure 13. The following is a list of useful detection, capture probes, and positive hybridization control probes. The detection, capture and positive hybridization control probes are also listed in the Sequence Listing as SEQ. ID. Nos. 32-43.

	SEQ. ID. No.	Capture Probes:
30	32	CAP235: TGT ATT AAC TGT CAA AAG CCA BIOTIN
	33	CAP250: TGT ATT AAC TGT CAA AAG CCA AAA AAA
		BIOTIN

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	34	CAP 253: TGT ATT AAC TGT CAA AAG CCA AAA AAA
		AAA A BIOTIN
	35	CAP265: GTA GAG AAA CCC AGC TGT AAA AAA
		BIOTIN
5	36	CAP267: GTG CCT GCG GTG CCA GAA AAA AAA
		BIOTIN
	SEO ID No	Detection Probes

	SEO. JD. No.	<u>Detection Probes</u> :
	37	DET59: GAC AGT ATT GGA ACT TAC AG
10	38	DET98: TTA GAA TGT GTG TAC TGC AAG NH2
	39	DET255: CAA CAG TTA CTG CGA CGT GAG NH2
	40	DET256: TTA CTG CGA CGT GAG GT NH2
	41	DET260: GTA TAT TGC AAG ACA GTA NH2
15	SEO. ID. No.	Positive Hybridization Control Probes:
	42	PHC271: TGT CTT GCA ATA TAC AAA AA BIOTIN
	43	PHC272: CTC ACG TCG CAG TAA AAA AAA BIOTIN

probes using 4 different detection probes. The capture probes were hybridized to the 3SR product at the temperature optima for 30 minutes in 0.1% PVP, 2X SSC and 8 pmol capture probe. The reaction was applied to the microwell and allowed to incubate at room temperature for 20 minutes. The microwell was washed 3 times in 2X SSC, 0.05% Tween 20 and 0.01% Thimersol. The detection probe was added to the microwell and hybridized at room temperature for 30 minutes. The well was again washed 3 times and developed for 15 minutes. The reaction was stopped and read at A450. The performance of the capture probes on the plate assay could be increased by the addition of adenine residues on the end of the oligos closest to the well (data not shown). Different bases were targeted (G, C, A, and T). T was not chosen because most mRNA's are polyadenylated which would cause end hybridization. CAP 250 produces

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the highest signal when amplifying SiHa cells; however, CAP 250 only can capture two of the three spliced E6 RNA's. Several other capture probes were investigated and CAP 265 captures all three E6 transcripts. Each cell line splices E6 at different rates. CAP 265 was chosen because clinical specimens may be heterogenous in splicing E6.

Once the capture probe has been defined, selecting an enzyme-conjugated detection probe is undertaken. Figure 17 is a comparison of all the detection probes for HPV 16. DET 256 produces the highest absorbance values in the present assay. Two detection probes were synthesized for illustration. The first a 17mer and the second a 15mer to define the minimum number of bases needed for efficient hybridization. The minimum length a detector oligo can be is about 17 bases (figure 18). Please note that best results are achieved when the signal enzyme is conjugated to the oligonucleotide at the 3' end.

Various additives in the capture buffer were performed with little increase in the relative absorbance in the plate assay (figure 19). When these same additives were added to the detection buffer the signal was more than doubled (figure 20). This effect appears to be related to the length of the 3SR product. The longer the product the more pronounced the effect. Primers 96-91 produce a shorter 3SR product than 136-91 (figure 20). Including additives in the detection buffer increases background levels. A titration using glycerol reduces background levels. Figure 21 is an autoradiogram of additional primer set that amplify HPV 18 using HeLa RNA. Figure 22 demonstrates the performance of HPV 18 capture probes using a variety of detection probes. Figure 23 demonstrates the absorbance values of a co-amplification and co-capture of HPV 16 and HPV 18 using type specific detection probes. Best results were achieved in coamplification for HPV 16 and HPV 18 simultaneously utilizing primers 136-91 (HPV 16), 54-69 (HPV 18), CAP 265 (HPV 16), CAP 267 (HPV 18), and DET 256 (HPV 16), DET 60 (HPV 18) as shown in figure 24. The configuration of this assay is shown in figure 25.

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The Assay Format. Utilizing the reagents described hereinabove, the assay format of the present invention was devised to optimize the signal obtainable from specimens having low viral mRNA copy number. A fluid phase capture of sample target sequence complementary to a capture prove sequence is much more efficient than adsorbing directly onto a solid phase. In fact, in a typical sandwich configuration, it is not uncommon to capture only 1-3% of total available nucleic acid in the sample. This reduces sensitivity correspondingly by two orders of magnitude.

Since it is still necessary to separate nucleic acid complexes on a solid phase, the "capture" sample must be immobilized onto the solid phase before the detection probe is added. The present assay takes advantage of the extremely high binding constant for the interaction between biotin and streptavidin. The capture oligonucleotide is biotinylated through 3' or 5' terminal labeling by conventional techniques. It has been empirically determined for the probes studied to date that biotinylating the capture probe at the 3' terminus is more efficient in immobilizing the probe hybridized to sample target sequence.

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The solid phase is coated with streptavidin, so that when the hybridized capture-sample sequence complex is brought into contact with it, the reaction between streptavidin and biotin takes place. The solid phase is preferably the inner surface of microtiter tray wells, but any solid phase separation system known to the art is satisfactory including but not limited to polystyrene beads, magnetic microparticles, test strips of plastic or metal, dipsticks, columns packed with a variety of materials, etc. The fluid phase capture method of the present invention is expected to give enhanced results with solid supports made of plastic because of the especially low capture efficiencies with plastic supports in conventional assays.

Any signal-generating enzyme or other reporter or tracer system capable of being conjugated covalently or electrostatically to a oligonucleotide without hindering its hybridizing to a complementary

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sequence is contemplated in the present assay. Horseradish peroxidase is preferred, but alkaline phosphatase and synthetic fluorogenic and chromogenic molecule hydrolyzing enzymes may also be employed. Non-isotopic reporter/tracer systems are preferred over radioactive tracers because of environmental and stability considerations.

The kinetics of hybridization of various capture and detection probes will differ according to their thermodynamic characteristics, and some relatively insignificant amount of experimentation may be required to optimize the assay for probes of similar but not identical sequence disclosed herein for illustrative purposes.

Alternative Amplification Reaction Conditions

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Figure 26 compares amplification reactions performed using the standard 3SR reaction conditions (42°C) with amplification reactions performed at an elevated temperature (50°C). The assays used the primer sets 136-91 (HPV 16) and 54-69 (HPV 18) together and separately. The standard 3SR reaction conditions were 40 mM Tris-HCl, pH 8.1; 30 mM MgCl₂; 20 mM KCl; 10 mM dithiothreitol; 4 mM spermidine; 15 pmole each priming oligonucleotide; 1 mM dNTP's; 7 mM rNTP's; 30 units AMV reverse transcriptase; 2 units RNAse H; and 1000 units T7 RNA polymerase. The reaction was incubated for 1 hour at 42°C. The elevated temperature reaction conditions were 40 mM Tris acetate, pH 8.1; 30 mM Mg acetate; 10 mM dithiothreitol; 100 mM potassium glutamate, pH 8.1; 1 mM dNTP's; 6 mM rNTP's; 15% sorbitol; 30 units AMV reverse transcriptase; 2 units RNAse H; and 1000 units T7 RNA polymerase. The reaction was incubated for 1 hour at 50°C.

After incubating the amplification reactions, 1/10th of the amplification products were denatured in 90 μ l of 7.4% formaldehyde and 10X SSC in a 65°C water bath for 10 minutes and quick-chilled on ice for at least 1 minute. BA-85 nitrocellulose was pre-wetted with water and then with 10X SSC. The denatured amplification samples were applied to a slot blot apparatus containing the pre-wetted nitrocellulose and the samples

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were drawn onto the nitrocellulose using a vacuum. The filter was then baked for 45 minutes at 80°C and hybridized with a type-specific oligonucleotide specific for HPV 18 (DET59) or HPV 16 (DET98). The hybridization solution contains 6X SSC; 10X Denhardts; 10 mM Tris, pH 7.4; 0.2 mg/ml sheared salmon sperm DNA; and 1% SDS.

Figures 26 and 27 depict a comparison of the amplification yields of reactions performed at 50°C and at 42°C. In both figures, the amplification reactions in column 1 used the HPV 16 primers 136-91, the reactions in column 2 used the HPV 18 primers 54-69, and the reactions in column 3 used a combination of the HPV 16 and HPV 18 primers 136-91 and 54-69. The target sequence was a mixture of 5 amol each of SiHa cell (infected with HPV 16) and HeLa cell (infected with HPV 18) RNA. Rows 1-4 contained sorbitol concentrations of 15%, 10%, 5% and 0% respectively; row 5 was a minus template reaction using 15% sorbitol; row 6 was blank; and rows 7-11 contained sorbitol concentrations of 15%, 10%, 5% and 0% respectively. Rows 1-5 were incubated at 50°C and rows 7-11 were incubated at 42°C. The amplification products in figure 26 were probed with DET 98 which is specific for HPV 16. The amplification products in figure 27 were probed with DET 59 which is specific for HPV 18.

Figure 26 depicts that the bands were much stronger at the 15% and 15% sorbitol levels than at the 5% or 0% levels. These results demonstrate that the increased sorbitol concentrations protect the enzymes so that the reaction can be incubated at 50°C rather than 42°C. When the sorbitol concentration was dropped below 10% the enzymes were not thermally protected and denatured at elevated temperatures, resulting in the decreased level of amplification. Figures 26 and 27 demonstrate that the elevated temperature increased the level of amplification when compared to the 42°C reaction conditions. This was particularly evident when the target sequence was co-amplified using the mixed primer set, 136-91 (HPV 16) and 54-69 (HPV 18). The estimated level of amplification using the elevated temperature was 10 fold higher than the level of amplification using the 42°C reaction conditions.

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The foregoing detailed description has been provided for a better understanding of the invention only and no unnecessary limitation should be understood therefrom as some modifications will be apparent to those skilled in the art without deviating from the spirit and scope of the appended claims.

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SEQUENCE LISTING

	(1) GENERAL INFORMATION:
5	(i) APPLICANT: Janice T. Brown
	(ii) TITLE OF INVENTION: HUMAN PAPILLOMAVIRUS DETECTION ASSAY
10	(iii) NUMBER OF SEQUENCES:44
	(iv) CORRESPONDENCE ADDRESS
1 -	(A) ADDRESSEE: Baxter Diagnostics Inc.
15	(B) STREET: One Baxter Parkway, Building DP-3E
	(C) CITY: Deerfield
20	(D) STATE: Illinois
	(E) COUNTRY: USA
.	(F) ZIP: 60015
25	(v)COMPUTER READABLE FORM (A) MEDIUM TYPE: Floppy disk
30	(B) COMPUTER: Apple Macintosh
30	(C) OPERATING SYSTEM: Apple Macintosh System 7.0
	(D) SOFTWARE: Macintosh Text File
35	(vi)CURRENT APPLICATION DATA (A) APPLICATION NUMBER: N/A
	(B) FILING DATE: N/A
40	(C) CLASSIFICATION: N/A
45	(vii)PRIOR APPLICATION DATA (A) APPLICATION NUMBER: US 08/058,920
	(B) FILING DATE: May 6, 1993
50	(viii)ATTORNEY/AGENT INFORMATION (A) NAME: Mark Buonaiuto
	(B) REGISTRATION NUMBER: 31,593
55	(C) REFERENCE/DOCKET NUMBER: BA-4448

23

(ix)TELECOMMUNICATION INFORMATION

5

(A) TELEPHONE: 708/948-2537

(B) TELEFAX: 708/948-2642

	(2)	INFORMATION FOR SEQ ID NO: 1
	(i) SEQUENCE CHARACTERISTICS
5		(A) LENGTH: 570
		(B) TYPE: nucleic acid
10		(C) STRANDEDNESS: double
10		(D) TOPOLOGY: linear
	(i	ii) HYPOTHETICAL: no
15	(i	v) ANTI-SENSE: no
•	(v virus	i) ORIGINAL SOURCE: (A) ORGANISM: Papaoviridae, Human papilloma
20		(B) STRAIN: 16
	(i	x) FEATURE:
25	E6/E7 p	(A) NAME/KEY: Portion of viral genome coding for olypeptides.
30	(х М.,) PUBLICATION INFORMATION: (A) AUTHORS: Seedorf, K., Krammer, G., Durst,
	,	Suhai, S., and Rowekamp, W.
35	Sequenc	(B) TITLE: Human Papillomavirus Type 16 DNA e
		(C) JOURNAL: Virology
		(D) VOLUME: 145
40		(E) ISSUE:
		(F) PAGES: 181-185
45		(G) DATE: 1985
	(x	i) SEQUENCE DESCRIPTION: SEQ ID NO: 1
50	CGA 4	AC CAA AAG AGA ACT GCA ATG TTT CAG GAC CCA CAG GAG 6 is Gln Lys Arg Thr Ala Met Phe Gln Asp Pro Gln Glu
	Arg	5 10
	15	

	ATA	91	AAG											
	Pro Ile	Arg	Lys	Leu	Pro	Gln	Leu	Суѕ	Thr	Glu	Leu	Gln	Thr	Thr
5	30				20					25				
	CAT		ATA	ATA	TTA	GAA	TGT	GTG	TAC	TGC	AAG	CAA	CAG	TTA
10	His Leu	Asp	Ile	Ile	Leu	Glu	Суѕ	Val	Tyr	Cys	Lys	Gln	Gln	Leu
	45				35					40				
15		CGT 181	GAG	GTA	TAT	GAC	TTT	GCT	TTT	CGG	GAT	TTA	TGC	ATA
			Glu	Val	Tyr	Asp	Phe	Ala	Phe	Arg	Asp	Leu	Cys	Ile
20	60				50					55				
			GAT	GGG	AAT	CCA	TAT	ĢCT	GTA	TGT	GAT	AAA	TGT	TTA
25	Tyr	226 Arg	Asp	Gly	Asn	Pro	Tyr	Ala	Val	Cys	Asp	Lys	Cys	Leu
25	Lys 75				65					70				
			TCT	AAA	ATT	AGT	GAG	TAT	AGA	CAT	TAT	TGT	ТАТ	AGT
30		271 Tyr	Ser	Lys	Ile	Ser	Glu	Tyr	Arg	His	Tyr	Cys	Tyr	Ser
	Leu				80		:			85				
35	90						•							
	GAT 3	316	ACA											
	Tyr Asp	Gly	Thr	Thr		Glu	Gln	Gln	Tyr	Asn	Lys	Pro	Leu	Cys
40	105				95	-				100				
			ATT	AGG	TGT	ATT	AAC	TGT	CAA	AAG	CCA	CTG	TGT	ССТ
45	Leu	361 Leu	Ile	Arg	Cys	Ile	Asn	Cys	Gln	Lys	Pro	Leu	Cys	Pro
	Glu 120				110					115				
50			CAA	AGA	CAT	CTG	GAC	AAA	AAG	CAA	AGA	TTC	CAT	AAT
	Glu	Lys	Gln	Arg	His	Leu	Asp	Lys	Lys	Gln	Arg	Phe	His	Asn
55	Ile				125					130				

	TCA 451				•		GC AGA TC	
5	Ser	Arg rip	140	Arg Cys	145		s Alg Se.	L
	150							
10	AGA ACA	CGT AGA	GAA ACC	CAG CTG	TAATC A	TG CAT	GGA GAT	ACA
	Arg Thr	Arg Arg	Glu Thr 155	Gln Leu	M	let His	Gly Asp '	Thr 5
15	CCT ACA ACT 540	TTG CAT	GAA TAT	ATG TTA	GAT TTG	CAA CO	CA GAG AC	A
	Pro Thr Thr	Leu His	_	Met Leu	Asp Leu		o Glu Thi	r
	20		10		10	•		
20	GAT CTC	TAC TGT	TAT GAG	CAA TTA	AAT GAC	2		
25	Asp Leu	Tyr Cys	Tyr Glu 25	Gln Leu	Asn Asp 30			
23								
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		(D)	TOPOLOG	Y: linea	ar			
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40	(iv)	ANTI-	SENSE: 1	no				
	(vi)	ORIGII		CE: Papovavi:	ridae, H	Iuman pa	apilloma	
45	virus							
		(B) ST	RAIN: 18					
50	(vii	i) POSIT (A) CH	ION IN G					
	(ix)							_
55	E6/E7 poly			Portion (of viral	genome	e coding	IOI

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-	0		•	B)	TITL	E:	Nu	cleo	tide	Seq	uenc	e an	d		
5		para					An	alys	is o	f th	e Hu	man			
	Pap.	1110	mavi	rus			Ty	pe 1	8 Ge	nome	•				
10			(C)	JOUR	NAL:	Jo	urna	l of	Mol	ecul	ar B	iolo	gy	
			(1	D)	VOLU	ME:	19	3							
			()	E)	ISSU	E:									
15			(1	F)	PAGE	s:	59	9-60	8						
			((G)	DATE	:	19	87							
20															
		(x:			UENC								_		
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25	Met	Ala	Arg	Phe	Glu 5	Asp	Pro	Thr	Arg	Arg 10	Pro	Tyr	Lys	Leu	Pro 15
	GAT 90	CTG	TGC	ACG	GAA	CTG	AAC	ACT	TCA	CTG	CAA	GAC	ATA	GAA	ATA
30		Leu	Cys	Thr	Glu 20	Leu	Asn	Thr	Ser	Leu 25	Gln	Asp	Ile	Glu	Ile 30
	ACC 135	TGT	GTA	TAT	TGC	AAG	ACA	GTA	TTG	GAA	CTT	ACA	GAG	GTA	TTT
35		Cys	Val	Tyr	Cys 35	Lys	Thr	Val	Leu	Glu 40	Leu	Thr	Glu	Val	Phe 45
	GAA 180	TTT	GCA	TTT	AAA	GAT	TTA	TTT	GTG	GTG	TĄT	AGA	GAC	AGT	'ATA
40		Phe	Ala	Phe	Lys 50	Asp	Leu	Phe	Val	Val 55		Arg	Asp	Ser	Ile 60
	CCG 225	CAT	GCT	GCA	TGC	CAT	AAA	TGT	ATA	GAT	TTT	TAT	TCT	AGA	ATT
45		His	Ala	Ala	Cys 65	His	Lys	Cys	Ile	Asp 70	Phe	Tyr	Ser	Arg	Ile 75
	AGA 270	GAA	TTA	AGA	CAT	TAT	TCA	GAC	TCT	GTG	TAT	GGA	GAC	ACA	TTG
50		Glu	Leu ·	Arg	His 80	Tyr	Ser	Asp	Ser	Val 85	Tyr	Gly	Asp	Thr	Leu 90
	GAA 315	AAA	CTA	ACT	AAC	ACT	GGG	TTA	TAC	AAT	TTA	TTA	ATA	AGG	TGC
55	Glu	Lys	Leu	Thr	Asn	Thr	Gly	Leu	Tyr	Asn	Leu	Leu	Ile	Arg	Cys

	95 100 105
	CTG CGG TGC CAG AAA CCG TTG AAT CCA GCA GAA AAA CTT AGA CAG
5	Leu Arg Cys Gln Lys Pro Leu Asn Pro Ala Glu Lys Leu Arg His 110 115 120
	CTT AAT GAA AAA CGA CGA TTT CAC AAC ATA GCT GGG CAC TAT AGA
10	Leu Asn Glu Lys Arg Arg Phe His Asn Ile Ala Gly His Tyr Arg 125 130
	GGC CAG TGC CAT TCG TGC TGC AAC CGA GCA CGA CAG GAA CGA CTC 450
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	(D) TOPOLOGY: linear
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	DNA (iii) HYPOTHETICAL: no
40	(iv) ANTI-SENSE: no
	(vii) IMMEDIATE SOURCE:
45	(A) LIBRARY: DNA synthesizer
	<pre>(ix) FEATURE:</pre>
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3
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		(B) TYPE: nucleic acid					
10		(C) STRANDEDNESS: single					
		(D) TOPOLOGY: linear					
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		(iii) HYPOTHETICAL: no					
20		(iv) ANTI-SENSE: no					
20		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>					
25		(ix) FEATURE: (A) NAME/KEY: HPV19.					
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4					
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		(B) TYPE: nucleic acid					
40		(C) STRANDEDNESS: single					
		(D) TOPOLOGY: linear					
4 5	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic					
•		(iii) HYPOTHETICAL: no					
50		(iv) ANTI-SENSE: no					
		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>					
55		(ix) FEATURE.					

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	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20	(ii) MOLECULE TYPE:	
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25	(iii) HYPOTHETICAL: no	
	(iv) ANTI-SENSE: no	
30	(ix) FEATURE: (A) NAME/KEY: HPV29.	

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5	(2)	IN	FORMATI(ON FOR SEQ	ID NO:	7		
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10			(A)	LENGTH:	22			
			(B)	TYPE: nu	cleic a	cid		
4.5			(C)	STRANDEDN	ESS: s	ingle		
15			(D)	TOPOLOGY:	linea	r		
20	DNA			ULE TYPE: SCRIPTION:	Other	nucleic	acid,	synthetic
		(iii)	HYPOTI	HETICAL:	no			
		(iv)	ANTI-	SENSE: no				
25				IATE SOURCE BRARY: Di		hesizer		
30			FEATUI	RE: ME/KEY: HP	V32.			
		(xi)	SEQUE	NCE DESCRI	PTION:	SEQ ID	NO:7	
35	CACT	ICACTG	CAAGAC	ATAG AA			22	
	(2)	IN	FORMATIO	ON FOR SEQ	ID NO:	8		
40		(i)	SEQUE	NCE CHARAC	TERISTI	CS		
			(A)	LENGTH:	46			
			(B)	TYPE: nu	cleic a	cid		
45			(C)	STRANDEDN	ESS: s	ingle		
			(D)	TOPOLOGY:	linea	r		
50	DNA	(ii)	MOLECT (A) DE:	ULE TYPE: SCRIPTION:	Other	nucleic	acid,	synthetic
		(iii)	HYPOT	HETICAL:	no			
e e		/ i \	A NUTE -	CENCE				

	<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
5	(ix) FEATURE: (A) NAME/KEY: HPV48.
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8
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	(C) STRANDEDNESS: single
25	(D) TOPOLOGY: linear
23	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic DNA
30	(iii) HYPOTHETICAL: no
	(iv) ANTI-SENSE: no
35	<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
40	(ix) FEATURE: (A) NAME/KEY: HPV53.
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9
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55	(C) STRANDEDNESS: single

	(D) TOPOLOGY: li	near			
5	(ii) MOLECULE TYPE: (A) DESCRIPTION: Oth	ner nucleic acid, synthetic			
	(iii) HYPOTHETICAL: no				
10	(iv) ANTI-SENSE: no				
10	(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA s	ynthesizer			
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	(xi) SEQUENCE DESCRIPTION	N: SEQ ID NO:10			
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	(A) LENGTH: 21				
30	(B) TYPE: nuclei	c acid			
	(C) STRANDEDNESS:	single			
	(D) TOPOLOGY: li	near			
35	(ii) MOLECULE TYPE: (A) DESCRIPTION: Oth DNA	er nucleic acid, synthetic			
4.0	(iii) HYPOTHETICAL: no				
40	(iv) ANTI-SENSE: no				
45	(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA s	ynthesizer			
43	(ix) FEATURE: (A) NAME/KEY: HPV69.				
50	(xi) SEQUENCE DESCRIPTIO	N: SEQ ID NO:11			
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· 55	(2) INFORMATION FOR SEQ ID	NO:12			

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		(A) LENGTH: 23
5		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
10		(D) TOPOLOGY: linear
10	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
15		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
20		(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer
		(ix) FEATURE: (A) NAME/KEY: HPV73.
25		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12
	CAGT	PATGCA CAGAGCTGCA AAC 23
30	(2)	INFORMATION FOR SEQ ID NO:13
		(i) SEQUENCE CHARACTERISTICS
		(A) LENGTH: 23
35		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
40		(D) TOPOLOGY: linear
	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
45		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
50		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
55		(ix) FEATURE: (A) NAME/KEY: HPV74.

		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13
	GTTA	TGCACA GAGCTGCAAA CAA 23
5	(2)	INFORMATION FOR SEQ ID NO:14
		(i) SEQUENCE CHARACTERISTICS
10		(A) LENGTH: 20
		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
15		(D) TOPOLOGY: linear
20	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, syntheti
		(iii) HYPOTHETICAL: no
, 3 E		(iv) ANTI-SENSE: no
25		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
30		(ix) FEATURE: (A) NAME/KEY: HPV77.
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14
35	CAAG	CAACAG TTACTGCGAC 20
	(2)	INFORMATION FOR SEQ ID NO:15
40		(i) SEQUENCE CHARACTERISTICS
		(A) LENGTH: 20
		(B) TYPE: nucleic acid
4 5		(C) STRANDEDNESS: single
		(D) TOPOLOGY: linear
50	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
		(iii) HYPOTHETICAL: no
55		(iv) ANTI-CENCE: no

	(1		IATE SOURCE BRARY: DN		izer	
5	i)	.x) FEATU (A) NA	RE: ME/KEY: HPV	89.		
	(2	(i) SEQUE	NCE DESCRIP	TION: SE	Q ID NO:15	
10	AGCAACA	GTT ACTGCG	ACGT		20	
	(2)	INFORMATI	ON FOR SEQ	ID NO:16		
15	(:) SEQUE	NCE CHARACT	ERISTICS		
		(A)	LENGTH:	23		
		(B)	TYPE: nuc	leic acid	I	
20		(C)	STRANDEDNE	SS: sing	rle	
		(D)	TOPOLOGY:	linear		
25	DNA	i) MOLEC (A) DE	TULE TYPE:	Other nuc	eleic acid,	synthetic
30	(:	Lii) HYPOT	HETICAL:	no		
30	(:	Lv) ANTI-	SENSE: no			
35	7)		IATE SOURCE BRARY: DN		sizer	
33	(:	ix) FEATU (A) NA	RE: ME/KEY: HPV	90.		
40	(2	ki) SEQUE	NCE DESCRIP	TION: SE	Q ID NO:16	
40	GCACAG	AGCT GCAAAC	AACT ATA			23
45	(2)	INFORMATI	ON FOR SEQ	ID NO:17		
45	(:	i) SEQUE	NCE CHARACT	ERISTICS		
		(A)	LENGTH:	23		
50		(B)	TYPE: nuc	leic acid	ì	
		(C)	STRANDEDNE	SS: sing	,le	
55	٠	(D)	TOPOLOGY:	linear		

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	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
5	DIVA	(iii) HYPOTHETICAL: no
,		(iv) ANTI-SENSE: no
		(vii) IMMEDIATE SOURCE:
10		(A) LIBRARY: DNA synthesizer
		(ix) FEATURE: (A) NAME/KEY: HPV91.
15		(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17
	ACAGA	GCTGC AAACAACTAT ACA 23
20	(2)	INFORMATION FOR SEQ ID NO:18
		(i) SEQUENCE CHARACTERISTICS
25		(A) LENGTH: 51
25		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
30		(D) TOPOLOGY: linear
	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
35		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
40		(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer
		(ix) FEATURE: (A) NAME/KEY: HPV92.
45		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18
50	AATT: 51	PANTAC GACTCACTAT AGGGACTTTT CTTCAGGACA CAGTGGCTTT T
	(2)	INFORMATION FOR SEQ ID NO:19
	, - ,	(i) SEQUENCE CHARACTERISTICS
55		

		(A)	LENGTH: 50	
		(B)	TYPE: nucleic acid	
5		(C)	STRANDEDNESS: single	
		(D)	TOPOLOGY: linear	
10	(DNA		ECULE TYPE: DESCRIPTION: Other nucleic acid, syntheti	.c
	(iii) HYP	OTHETICAL: no	
15	(iv) ANT	I-SENSE: no	
	(EDIATE SOURCE: LIBRARY: DNA synthesizer	
20	(ix) FEA	TURE: NAME/KEY: HPV93.	
	(xi) SEQ	UENCE DESCRIPTION: SEQ ID NO:19	
25	AATTTA 50	ATAC GACT	CACTAT AGGGATTTGC TTTTCTTCAG GACACAGTGG	
	(2)	INFORMA	TION FOR SEQ ID NO:20	
30	(i) SEQ	UENCE CHARACTERISTICS	
		(A)	LENGTH: 50	
35		(B)	TYPE: nucleic acid	
		(C)	STRANDEDNESS: single	
40		(D)	TOPOLOGY: linear	
	(DNA		ECULE TYPE: DESCRIPTION: Other nucleic acid, syntheti	.c
45	(iii) HYP	OTHETICAL: no	
	(iv) ANT	I-SENSE: no	
50	(EDIATE SOURCE: LIBRARY: DNA synthesizer	
	(ix) FEA (A)	TURE: NAME/KEY: HPV94.	
55	(xi) SEO	UENCE DESCRIPTION: SEO ID NO:20	

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	AATTTAATAC GACTCACTAT AGGGATCTTT GCTTTTCTTC AGGACACAGT 50						
5	(2) INFORMATION FOR SEQ ID NO:21						
	(i) SEQUENCE CHARACTERISTICS						
10	(A) LENGTH: 50						
	(B) TYPE: nucleic acid						
15	(C) STRANDEDNESS: single						
13	(D) TOPOLOGY: linear						
20	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic DNA						
	(iii) HYPOTHETICAL: no						
	(iv) ANTI-SENSE: no						
25	<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>						
30	(ix) FEATURE: (A) NAME/KEY: HPV95.						
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21						
35	AATTTAATAC GACTCACTAT AGGGATGTCT TTGCTTTTCT TCAGGACACA 50						
	(2) INFORMATION FOR SEQ ID NO:22						
40	(i) SEQUENCE CHARACTERISTICS						
	(A) LENGTH: 50						
45	(B) TYPE: nucleic acid						
43	(C) STRANDEDNESS: single						
	(D) TOPOLOGY: linear						
50	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic DNA						

no

(iii) HYPOTHETICAL:

	(iv) ANTI-SENSE: no	
5	<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>	
)	(ix) FEATURE: (A) NAME/KEY: HPV96.	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22	
10	AATTTAATAC GACTCACTAT AGGGAGATGT CTTTGCTTTT CTTCAGGACA 50	
15	(2) INFORMATION FOR SEQ ID NO:23	
	(i) SEQUENCE CHARACTERISTICS	
20	(A) LENGTH: 23	
20	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
25	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, syntheti DNA	.c
30	(iii) HYPOTHETICAL: no	
	(iv) ANTI-SENSE: no	
35	(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer	
40	(ix) FEATURE: (A) NAME/KEY: HPV101.	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23	
	AGAGCTGCAA ACAACTATAC ATG 23	
4 5	(2) INFORMATION FOR SEQ ID NO:24	
	(i) SEQUENCE CHARACTERISTICS	
50	(A) LENGTH: 49	
	(B) TYPE: nucleic acid	
55	(C) STRANDEDNESS: single	

		(D) TOPOLOGY: linear
5	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
		(iii) HYPOTHETICAL: no
1.0		(iv) ANTI-SENSE: no
10		(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer
15		(ix) FEATURE: (A) NAME/KEY: HPV106.
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24
20	AATT 49	PAATAC GACTCACTAT AGGGATTCAT GCAATGTAGG TGTATCTCC
	(2)	INFORMATION FOR SEQ ID NO:25
25		(i) SEQUENCE CHARACTERISTICS
		(A) LENGTH: 49
30		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
		(D) TOPOLOGY: linear
35	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
		(iii) HYPOTHETICAL: no
40		(iv) ANTI-SENSE: no
45		(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer
45		(ix) FEATURE: (A) NAME/KEY: HPV107.
50		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25
30	AATT 49	PANTAC GACTCACTAT AGGGATATTC ATGCAATGTA GGTGTATCT
55	(2)	INFORMATION FOR SEQ ID NO:26

		(i)	SEQUE	NCE CHARA	CTERIS	rics		
5			(A)	LENGTH:	23			
5			(B)	TYPE: n	ucleic	acid		
			(C)	STRANDED	NESS:	single		
10			(D)	TOPOLOGY	: line	ear		
1.5	DNA	(ii)		ULE TYPE: SCRIPTION		r nucleic	acid,	synthetic
15		(iii)	HYPOT	HETICAL:	no			
		(iv)	ANTI-	SENSE: n	10			
20		(vii)		IATE SOUF BRARY:		nthesizer		
25		(ix)	FEATU (A) NA	RE: ME/KEY: H	IPV118.			
		(xi)	SEQUE	NCE DESCR	RIPTION	: SEQ ID	NO:26	
	AGCT	GCAAAC	C AACTAT.	ACAT GAT			2	3
30	(2)	II	NFORMATI	ON FOR SE	EQ ID NO	0:27		
		(i)	SEQUE	NCE CHARA	CTERIS	TICS		
35			(A)	LENGTH:	4	9		
			(B)	TYPE: r	nucleic	acid		
40			(C)	STRANDEL	NESS:	single		
			(D)	TOPOLOGY	: line	ear		
45	DNA	(ii)		ULE TYPE: SCRIPTION		r nucleic	acid,	synthetic
		(iii)	HYPOT	HETICAL:	no			
E O		(iv)	ANTI-	SENSE: r	10			
50			IMMED	IATE SOUF	RCE:	nthesizer		

	bind:	ing si		AME/KEY: HPV120. Phage T/ RNA polymerase 5'end, followed by HPV-16/18 sequence.
_		(xi)	SEQU	ENCE DESCRIPTION: SEQ ID NO:27
5	AATT' 49	ГААТАС	GACTC	ACTAT AGGGATGCAA TGTAGGTGTA TCTCCATGC
10	(2)	IN	IFORMAT:	ION FOR SEQ ID NO:28
		(i)	SEQU	ENCE CHARACTERISTICS
1 F			(A)	LENGTH: 48
15			(B)	TYPE: nucleic acid
			(C)	STRANDEDNESS: single
20			(D)	TOPOLOGY: linear
	DNA	(ii)	MOLE (A) D	CULE TYPE: ESCRIPTION: Other nucleic acid, synthetic
25		(iii)	НҮРО	THETICAL: no
		(iv)	ANTI	-SENSE: no
30		(vii)		DIATE SOURCE: IBRARY: DNA synthesizer
35		(ix)	FEAT	URE: AME/KEY: HPV129.
33		(xi)	SEQU	ENCE DESCRIPTION: SEQ ID NO:28
	AATT	TAATAC	GACTC	ACTAT AGGGAAATGT AGGTGTATCT GGATGCAT 48
40	(2)	IN	IFORMAT	ION FOR SEQ ID NO: 29
		(i)	SEQU	ENCE CHARACTERISTICS
45			(A)	LENGTH: 23
			(B)	TYPE: nucleic acid
50			(C)	STRANDEDNESS: single
J 0			(D)	TOPOLOGY: linear
55	מארז	(ii)	MOLE (A) D	CULE TYPE: ESCRIPTION: Other nucleic acid, synthetic

	(i	ii) HYPO	THETICAL:	no		
-	(i	v) ANTI-	-SENSE: .no			
5	(v		DIATE SOURCE IBRARY: DA		esizer	
10	(i	x) FEAT	JRE: AME/KEY: HP\	7131.		
	(×	i) SEQUI	ENCE DESCRII	PTION: S	SEQ ID NO:	29
15	AAACAAC	TAT ACATG	ATATA ATA		23	
	(2)	INFORMAT	ION FOR SEQ	ID NO:30)	
20	(i	.) SEQUI	ENCE CHARACT	TERISTICS	3	
20		(A)	LENGTH:	49		
		(B)	TYPE: nuc	cleic aci	ld	
25		(C)	STRANDEDNI	ESS: sir	ngle	
		(D)	TOPOLOGY:	linear		,
30	(i DNA		CULE TYPE: ESCRIPTION:	Other nu	ıcleic acid	, synthetic
	(i	.ii) HYPO	THETICAL:	no		
35	(i	.v) ANTI	-SENSE: no			
	(5		DIATE SOURCI IBRARY: DI		esizer	
40		x) FEAT (A) No r site at	URE: AME/KEY: HPV 5'end, follo	7136. Phowed by F	nage T7 RNA HPV-16/18 s	polymerase
			ENCE DESCRII			
45		_	ACTAT AGGGAA			
50	(2)	INFORMAT	ION FOR SEQ	ID NO:3	L	
	i)	.) SEQU	ENCE CHARACT	reristic:	3	
55		(A)	LENGTH:	49		

		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
5		(D) TOPOLOGY: linear
	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
10		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
15		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
20		(ix) FEATURE: (A) NAME/KEY: HPV137.
20		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31
25	AATT	FAATAC GACTCACTAT AGGGATGTAG GTGTATCTCC ATGCATGAT
	(2)	INFORMATION FOR SEQ ID NO:32
2.0		(i) SEQUENCE CHARACTERISTICS
30		(A) LENGTH: 21
		(B) TYPE: nucleic acid
35		(C) STRANDEDNESS: single
		(D) TOPOLOGY: linear
40	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
		(iii) HYPOTHETICAL: no
45		(iv) ANTI-SENSE: no
		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
50		(ix) FEATURE: (A) NAME/KEY: CAP245.
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32
EE	mcm ₂	መጥአ እርጥ ርጥር እ እ እ እርርር እ 21

	(2)	IN	FORMATIO	ON FOR S	EQ ID NO	0:33		
5		(i)	SEQUE	NCE CHAR	ACTERIST	rics		
			(A)	LENGTH:		27		
10			(B)	TYPE:	nucleic	acid		
10			(C)	STRANDE	DNESS:	single		
			(D)	TOPOLOG	Y: line	ear		
15	DNA	(ii)		ULE TYPE SCRIPTIO		nucleic	acid,	synthetic
20		(iii)	нүрот	HETICAL:	no			
20		(iv)	ANTI-	SENSE:	no			
25		(vii)		IATE SOU BRARY:		nthesizer		
23		(ix)	FEATU (A) NAI	RE: ME/KEY:	CAP250.			
30		(xi)	SEQUE	NCE DESC	RIPTION	SEQ ID	NO:33	
	TGTA'	TTAACT	GTCAAA	AGCC AAA	AAAA	27		
35	(2)			ON FOR S				
		(i)		NCE CHAR				
4.0				LENGTH:		31		
40				TYPE:				
	·			STRANDE		-		
45		(ii)	MOLEC	ULE TYPE	l :		agi d	synthetic
	DNA		(A) DE	SCRIPTIC	in: Other	nucleic	aciu,	synchecic
50		(iii)	НҮРОТ	HETICAL:	no			
		(iv)	ANTI-	SENSE:	no			
EE		(vii)		IATE SOU		nthesizer		

		(ix)	FEAT	TURE: JAME/KEY:	CAP253.				
5		(xi)	SEQU	JENCE DESC	RIPTION	SEQ	ID	NO:34	
	TGTA	TAACT	GTCA	AAGCC AAA	АААААА	A	31		
10	(2)	IN	FORMAT	TION FOR S	EQ ID NO	0:35			
		(i)	SEQU	JENCE CHAR	ACTERIS:	rics			
15			(A)	LENGTH:	2	24			
13			(B)	TYPE:	nucleic	acid			
			(C) .	STRANDE	DNESS:	single	e		
20			(D)	TOPOLOG	Y: line	ear			
	DNA	(ii)		ECULE TYPE DESCRIPTIO		r nucle	eic	acid,	synthetic
25		(iii)	НҮРС	OTHETICAL:	no				
		(iv)	ANT	I-SENSE:	no				
30		(vii)		EDIATE SOU LIBRARY:		nthesi	zer		
2.5		(ix)	FEAT	TURE: NAME/KEY:	CAP265.				
35		(xi)	SEQ	JENCE DESC	RIPTION	: SEQ	ID	NO:35	
	GTAG	AGAAAC	CCAG	CTGTAA AAA	.A		24	i	
40	(2)	IN	IFORMA'	TION FOR S	EQ ID N	D:36			
		(i)	SEQ	UENCE CHAR	ACTERIS	rics		•	
45			(A)	LENGTH:	24				
			(B)	TYPE:	nucleic	acid			
			(C)	STRANDE	DNESS:	singl	e		
50			(D)	TOPOLOG	Y: lin	ear			
55	DNA	(ii)	MOL:	ECULE TYPE DESCRIPTIO	E: ON: Othe	r nucl	eic	acid,	synthetic

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		(iii)	НУР	OTHETICA	AL:	no			
-		(iv)	ANT	'I-SENSE	no				
5				EDIATE S LIBRARY:			thesiz	er	
10				TURE: NAME/KE	r: CA	P267.			
		(xi)	SEÇ	UENCE DI	ESCRI	PTION:	SEQ	ID NO:36	
15	GTGCC'	TGCGG	TGCC	AGAAAA /	AAAA			24	
	(2)	IN	FORMA	TION FOR	R SEQ	ID NO	:37		
0.0		(i)	SEC	UENCE CI	HARAC'	TERIST	ics		
20			(A)	LENG'	CH:	20			
			(B)	TYPE	: nu	cleic	acid		
25			(C)	STRAI	NDEDN	ESS:	single	9	
			(D)	TOPO	LOGY:	line	ar		
30	DNA	(ii)		ECULE TO		Other	nucle	eic acid,	synthetic
		(iii)	нүн	POTHETIC	AL:	no			
35		(iv)	ANT	I-SENSE	: no				
				MEDIATE : LIBRARY			thesiz	zer	
40				ATURE:	Y: DE	т59.			

	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:37
	GACAG'	TATTG GAACTTACAG 20
5	(2)	INFORMATION FOR SEQ ID NO:38
		(i) SEQUENCE CHARACTERISTICS
10		(A) LENGTH: 21
		(B) TYPE: nucleic acid
15		(C) STRANDEDNESS: single
13		(D) TOPOLOGY: linear
20	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
25		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
30		(ix) FEATURE: (A) NAME/KEY: DET98.
		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38
35	TTAGA	ATGTG TGTACTGCAA G 21
	(2)	INFORMATION FOR SEQ ID NO:39
40		(i) SEQUENCE CHARACTERISTICS
10		(A) LENGTH: 21
		(B) TYPE: nucleic acid
45		(C) STRANDEDNESS: single
		(D) TOPOLOGY: linear
50	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
		(iii) HYPOTHETICAL: no
55		(iv) ANTI-SENSE: no

		(vii)		DIATE SO		synt	hesiz	er	
5.		(ix)	FEAT	URE: AME/KEY:	DET25	55.			
		(xi)	SEQU	ENCE DES	CRIPT	ON:	SEQ :	ID NO:39	
10	CAAC	AGTTAC	TGCGA	CGTGA G		2	1		
	(2)	IN	FORMAT	ION FOR	SEQ II	NO:	40		
15		(i)	SEQU	ENCE CHA	RACTE	RISTI	cs		
			(A)	LENGTH	:	17			
20			(B)	TYPE:	nucle	eic a	cid		
20			(C)	STRAND	EDNESS	S: s	ingle		
			(D)	TOPOLO	GY:]	inea	r ·		
25	DNA			CULE TYP		her	nucle:	ic acid,	synthetic
		(iii)	НҮРО	THETICAL	: no	>			
30		(iv)	ANTI	-SENSE:	no				
35		(vii)		DIATE SO		synt	hesize	er	
33			FEAT	URE: AME/KEY:	DET 2	256.			
40		(xi)	SEQU	ENCE DES	CRIPTI	ON:	SEQ :	ID NO:40	
-	TTAC	rgcgac	GTGAG	GT		17			
45	(2)	IN	FORMAT	ION FOR	SEQ II	NO:	41		
40		(i)	SEQU	ENCE CHA	RACTE	RISTI	cs		
			(A)	LENGTH	:	18			
50			(B)	TYPE:	nucle	eic a	cid		•
			(C)	STRAND	EDNESS	S: s	ingle		
55			(D)	TOPOLO	GY:]	linea	r		

	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
5		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
10		<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>
		(ix) FEATURE: (A) NAME/KEY: DET260.
15		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41
	GTAT	ATTGCA AGACAGTA 18
20	(2)	INFORMATION FOR SEQ ID NO:42
		(i) SEQUENCE CHARACTERISTICS
0.5		(A) LENGTH: 20
25		(B) TYPE: nucleic acid
		(C) STRANDEDNESS: single
30		(D) TOPOLOGY: linear
	DNA	(ii) MOLECULE TYPE: (A) DESCRIPTION: Other nucleic acid, synthetic
35		(iii) HYPOTHETICAL: no
		(iv) ANTI-SENSE: no
40		(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer
45		(ix) FEATURE: (A) NAME/KEY: PHC271.
40		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42
	TGTC'	TTGCAA TATACAAAAA 20
50	(2)	INFORMATION FOR SEQ ID NO:43
		(i) SEQUENCE CHARACTERISTICS
55		(A) LENCTH• 21

		(B)	TYPE:	nucleic	acid						
_		(C)	STRAND	EDNESS:	single						
5		(D)	TOPOLO	GY: line	ear						
10	(ii)		CULE TYP ESCRIPTI		r nucleic	acid,	synthetic				
	(iii	і) нуро	THETICAL	: no							
4.5	(iv)	ANTI	-SENSE:	no							
15	(vii	<pre>(vii) IMMEDIATE SOURCE: (A) LIBRARY: DNA synthesizer</pre>									
20	(ix)	FEAT		PHC272.							
	(xi)	SEQU	ENCE DES	CRIPTION	: SEQ ID	NO:43					
25	CTCACGTC	GC AGTAA	AAAAA A		21						
	(2)	INFORMAT	ION FOR	SEQ ID N	0:44						
30	(i)	SEQU	ENCE CHA	RACTERIS	TICS						
30		(A)	LENGTH	I:	25						
		(B)	TYPE:	nucleic	acid						
35		(C)	STRANI	EDNESS:	single						
		(D)	TOPOLO	GY: lin	ear						
	(xi) SEQU	ENCE DES	CRIPTION	: SEQ ID	NO:44					
40	AATTTAAT	AC GACTO	ACTAT AG	GGA	25						

31

I claim:

1	1. An assay of a patient specimen suspected of containing messenger
2	RNA encoded by at least one type of HPV associated with cervical
3	dysplasia, malignant cells, or pre-malignant cells comprising
4	(1) subjecting said specimen to nucleic acid amplification
5	by self sustained sequence replication utilizing two primers
6	separated by at least ten nucleotides, at least one such primer
7	containing a transcriptional promoter,
8	annealing the first said primer to its complementary
9	sequence on a target region of said messenger RNA, extending the 3'
10	end of said primer by action of a strand-extending polymerase in the
11	presence of cofactors and nucleotide triphosphates,
12	digesting the RNA strand of the nascent RNA/DNA
13	duplex with an enzyme RNAse H activity,
14	annealing the second said primer to its complementary
15	sequence on the resultant single stranded cDNA, primer extending
16	the 3' end of the primer by action of a strand-extending polymerase,
17	transcribing the double stranded DNA with a
18	transcriptase in the presence of nucleoside triphosphates, and
19	repeating the amplification utilizing the newly
20	synthesized transcripts as new targets,
21	(2) hybridizing in solution amplified messenger RNA to a
22	free biotinylated reagent capture probe have a sequence
23	complementary to a first segment of the amplified RNA to form a
24	reagent capture complex,
25	(3) attaching said capture complex to a solid phase by
26	reaction of the biotin residues of said capture probe with
27	streptavidin covalently bound to the surface of said phase,
28	(4) washing the bound capture complex to remove
2 9	unbound and unreacted reagents,
30	(5) hybridizing a virus type-specific reporter-conjugated

detection probe having a sequence complementary to a second

32	segment of the amplified RNA not overlapping the sequence of the
33	first such RNA segment to form a solid phase-bound capture probe-
34	target sequence-detection probe complex,
35	(6) washing the complex to remove unhybridized
36	detection probe, and
37	(7) adding a fluorogenic or chromogenic enzyme substrate
38	and reacting the conjugated enzyme to produce a detectable
39	fluorophor or chromogen.
1	2. An assay for detecting HPV in a cervical specimen associated with
2	cervical dysplasia or premalignant or malignant cells comprising
3	(1) amplifying target HPV messenger RNA encoding
4	sequences contained in the viral E6/E7 region which is contained in
5	said specimen by self sustained sequence replication,
6	(2) capturing said amplified messenger sequences by fluid
7	hybridization with a biotinylated capture probe having a sequence
8	complementary thereto,
9	(3) reacting said hybridized capture prove with a
10	streptavidin coated solid phase,
11	(4) washing to remove unbound hybridized capture probe,
12	(5) hybridizing a detection probe to said target sequence,
13	(6) washing said solid phase, and
14	(7) detecting the detecting probe.
1	3. An assay for detecting HPV in a cervical specimen associated with
2	cervical dysplasia or premalignant or malignant cells comprising
3	(1) coamplifying a plurality of oncogenic HPV type
4	messenger RNAs contained in said specimen and having sequences
5	encoding the respective E6/E7 genes of the HPV types or portions
6	thereof,

7	(2) capturing said amplified messenger sequences by fluid
8	hybridization with a biotinylated capture probe having a sequence
9	complementary thereto,
10	(3) reacting said hybridized capture probe with a
11	streptavidin coated solid phase,
12	(4) washing to remove unbound hybridized capture probe,
13	(5) hybridizing a detection probe to said target sequence,
14	(6) washing said solid phase, and
15	(7) detecting the detecting probe.

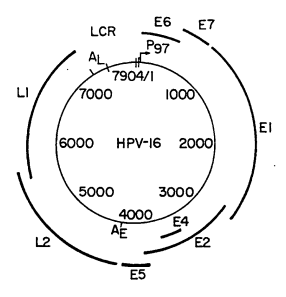
- 1 4. The assay of claims 1, 2, or 3 wherein said capture probes are selected
- 2 from the group consisting of CAP245, CAP250, CAP253, CAP265 and
- 3 CAP267.
- 1 5. The assay of claim 1 wherein the human papillomavirus-16 primers
- , 2 for self sustained sequence replication are selected from the group of
- 3 primer pairs consisting of HPV 16: 120-29, 120-90; 15-19, 15-20, 15-77, 15-53,
- 4 15-89, 15-29; 129-29, 129-74, 129-73, 129-118, 129-130, 129-131; 136-91, 136-29,
- 5 136-90, 136-74, 136-73, 136-130; 137-29, 137-90, 137-74, 137-73, 137-118; 93-73;
- 6 93-91; 85-77; 95-101, 95-91; 96-91, 96-73; 136-131; 94-91.
- 1 6. The assay of claims 1, 2, or 3 wherein said detection probes are
- 2 selected from the group consisting of DET256, DET255, DET98 and DET260.
- 1 7. Primer pairs for self sustained sequence amplification of the E6/E7
- 2 region of HPV-16 associated with cervical dysplasia or premalignant or
- 3 malignant cervical cells consisting of: 15-19, 15-20, 15-77, 15-53, 15-89, 15-29;
- 4 136-91, 136-29, 136-90, 136-74, 136-73, 136-130, 136-131, 136-118; 96-91, 96-73;
- 5 and 94-91.
- 1 8. Capture probes for capturing amplified RNA target sequences of the
- 2 HPV E6/E7 region consisting of CAP265 and CAP267.

- 1 9. Detection probes hybridizing to the E6/E7 region of HPV consisting
- 2 of enzyme-conjugated probes having the sequence of DET256, DET255,
- 3 DET98 and DET260.
- 1 10. Primer pairs for self sustained sequence amplification of the E6/E7
- 2 region of HPV-18 associated with cervical dysplasia or premalignant or
- 3 malignant cervical cells consisting of: 54-69, 54-70, 54-32.
- 1 11. The assay of claim 1 wherein the HPV-18 primers for self sustained
- 2 sequence replication are selected from the group of primer pairs consisting
- 3 of: 54-32, 54-69, 54-70; 48-32; 214-69, 214-244, 214-214, 214-70.
- 1 12. A kit for detection of HPV associated with cervical dysplasia,
- 2 premalignant or malignant cervical cells comprising any of the primer
- 3 pairs of claims 7 or 10, any of the capture probes of claim 8, and any of the
- 4 detection probes of claim 9.
- 1 13. The assay of claim 1 wherein said nucleic acid amplification by self
- 2 sustained sequence replication is performed at an elevated temperature of
- 3 about 50°C in the presence of a thermal protection agent.
- 1 14. The assay of claim 2 wherein said amplifying of said target RNA is
- 2 performed at an elevated temperature of about 50°C in the presence of a
- 3 thermal protection agent.
- 1 15. The assay of claim 3 wherein said coamplifying of said plurality of
- 2 RNAs is performed at an elevated temperature of about 50°C in the
- 3 presence of a thermal protection agent.

- 1 16. The assay of claim 1 wherein said patient sample is suspected of
- 2 containing messenger RNA encoded by the E6/E7 splice region of human
- 3 papillomavirus 16 or 18.
- 1 17. The assay of claim 2 wherein said viral E6/E7 region is from HPV 16
- 2 or 18.
- 1 18. The assay of claim 3 wherein said sequences encoding the E6/E7
- 2 genes are specific for the E6/E7 splice region of HPV 16 or 18.

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Fig.1



		T					R	T	A	ATG M	TTT F	CAG Q	GAC D	CCA P	CAG Q	GAG E	CG R	126
			CCC P	AGA R	AAG K	TTA L	CCA	HPV9 CAG Q	ATT	TGC C	ACA T	GAG E	L	u		ACT T	AT \	_171
_	-4-	A	CAT H	GAT D	ATA I	ATA I	TTA L	GAA E	TGT C	GTG V	TAC Y	TGC C	AAG K	256 — Caa Q	CAG Q	TTA L	CT L	_216
	\	G	CGA R	CGT R			Y	GAC D	TTT F	GCT A	TTT F	CGG R	GAT D	TTA L	TGC C	ATA I	GT V	261
		A		AGA R		GGG G	NH AAT N	CCA P	TAT Y	GCT A	GTA V	TGT C	GAT D	AAA K	TGT C	TTA L	AA K	306
		G	TTT F		TCT S	AAA K	ATT	AGT S	GAG E	TAT Y	AGA R	CAT H	TAT Y	TGT C	TAT Y	AGT S	TT L	351
		G	TAT Y			ACA T	TTA L	GAA E	CAG	CAA Q	TAC Y	AAC N	AAA K	CCG P	TTG L	TGT C	GA D	396
		т	TTG L		ATT I	AGC R	TGT C	ATT	AAC N	TGT C	CAA Q	AAG K	CCA P	CTG L	TGT C	CCT P	GA E	441
		A	GA#	AAG K	CAA Q	AGA R	CAT H	CTG L	GAC D	: AAA K	AAG K	CAA Q	AG#	TTC F	CAT H	AAT N	AT I	486
		A	AG(G GGT	CGG	TG6 W	ACC T	GGT G	CG#	C	M	5	TG1 C	TGC C	AG R	TCA S	TC S	531
		· A	CAF AG/ R	265 A AC/ T	A C <u>G</u> T	r AG/	A GA/	A ACC	CAC		136- TA/			•				565
	_	-~	AT(G CAT	r gg/	A GA	T AC	A CCT	r aca	A TITO	\H	E	A TA		BIOTI 3 TTA L		TTG L	606
Į	\		CA	A CC	A GAI	G AC	A AC	r ga ⁻	T CTI	C TAI	PBS CTG C	TAT	ΓGA E	G CA	A TT/	AAT N	r gac D	651

	GCG A		TTT F HPV8	E						CCC P	TAC Y	AAG K	CTA L	CCT P	162
D	CTG L 260—	C		GAA	CTG L	AAC N		TCA S	CTG L	CAA Q	GAC D	ATA I	GAA E	ATA I	207
			TAT Y	TGC C	AAG K	ACA T				CTT L		GAG E		TTT F	252
GAA E		GCA A	TTT F	AAA K	GAT D	TTA L		GTG V	GTG V	TAT Y	AGA R	GAC D	AGT S	ATA Į	297
	CAT H		GCA A	TGC C	CAT H	AAA K	TGT C	ATA I	GAT D	TTT F	TAT Y	TCT S	AGA R	ATT I	342
	GAA E				TAT Y		GAC D	TCT S	GTG V	TAT Y	G	GAC D 267-	T	TTG L	387
		CTA L		AAC N	ACT T	GGG G HPV	L		AAT N	TTA L	TTA	ATA		TGC C	432
 CTG L	CGG R	TGC C	CAG Q	AAA K	P	TTG L	AAT	P	GCA A	GAA E	AAA K	<u>CTT</u> L	AGA R	CAC 1	477
 CTT L				CGA R				AAC		GCT A	GGG G	CAC H	TAT Y	AGA R	522
GGC G					TGC C		AAC N	CGA R		CGA R	CAG Q	ĠAA E	CGA R	CTC L	567
CAA Q	CGA R	CGC R	AGA R	GAA E	ACA T	CAA Q	GTA V	TAA	TAT Y	TAA *					600

Fig.4	75 LSI217 2500 H2µ T 7 2500 8.6 BUFFER IO/3 PAULS×T	IOAM SI HO 6-7µg/\ IO%DMSO 15% SORB.				
129	9 1 29 74, u. 73	90 = 				
	3 8 - 9 29	90				
120	74 73 73 73 73 74 73 74 74 75 75 75 75 75 75 75 75 75 75 75 75 75	2 130 137 3				
136	91 29 74 75	30				
137	9 29 74 7,73 64 1,73 13 1,8	90 d				
106	74 73 74 73	90 150				
	91 29 74 73 131 118 150FM 10	90				

16100FM 134-b 136-90 136-74 136-73 HC-144 98 2PM 0.193 0.532 0.584 2.932 0.093

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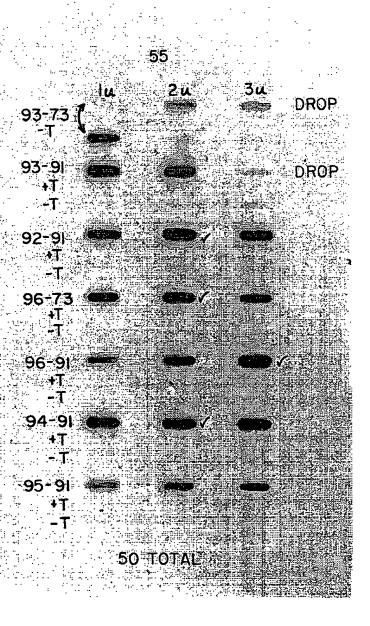


Fig. 6

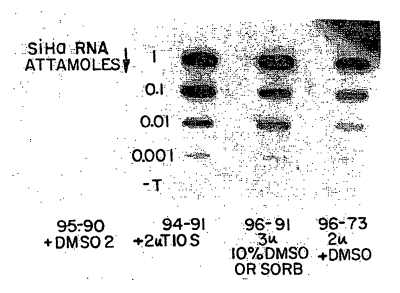


Fig. 7

TEMP AT 45°C 18 BB 33 O/N

TOTAL
RNA

32-48

IO⁴ IO-OIFM IO³ IO²

SOLN
TOTAL
RNA

IFW 35R IAM-35R -T
HELA

2hr 3SR

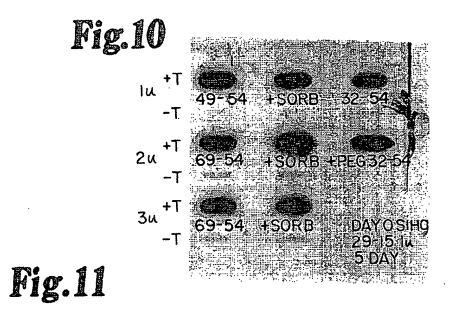
Fig. 8

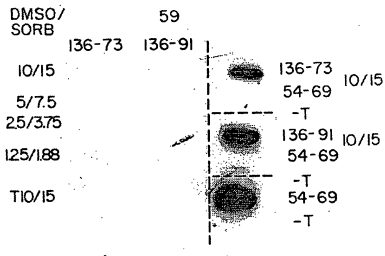
70°C 1.5 HR. **BB51** HELA 20 UNIT I.O UNIT 3.0 UNIT 20*uq* 32-54 #1 **-T** 32-54 #1 15 DNASE 32-54 #1 15, DNASE 32-54#2 **-T** 32-54#2 **DNAse** -T 32-54 Sind HELA PRIMERS 3.0m 1.0m 2.0m **DNASE** SiHa 1.0m 3.0m 2.0M 29-I5S **DNASE**

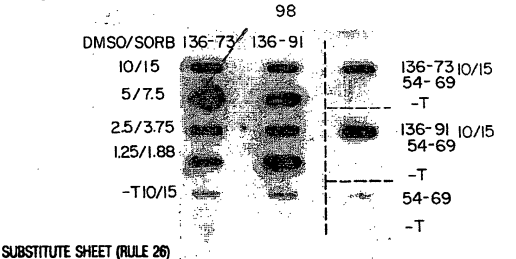
Fig. 9

PCT/US94/05085

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9/20 **Fig. 12**

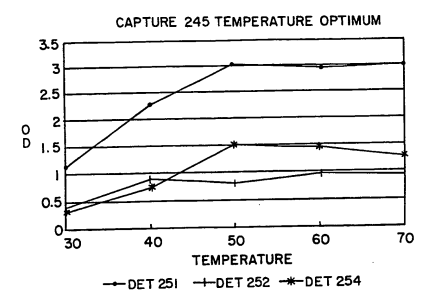
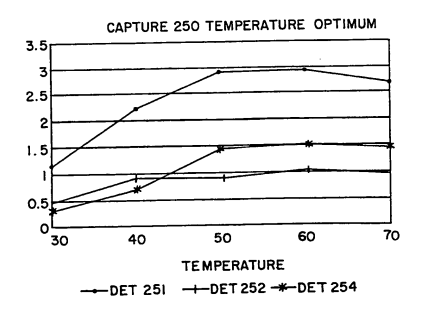


Fig. 13



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PCT/US94/05085

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Fig. 14

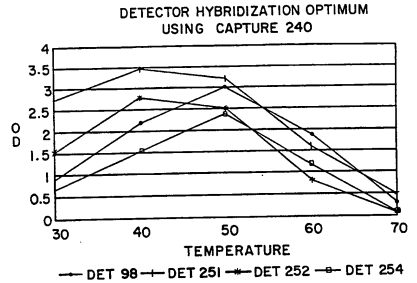
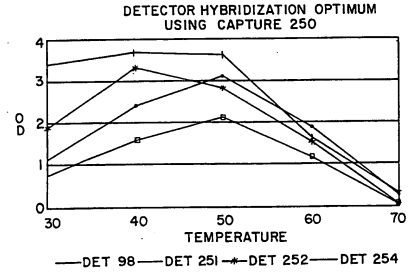


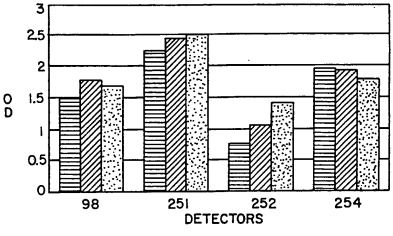
Fig. 15



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Fig. 16

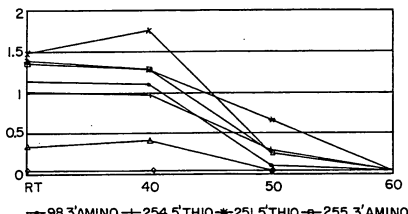
HPV CAPTURES AND DETECTORS



CAPTURES = 245 250 250 253

Fig. 17

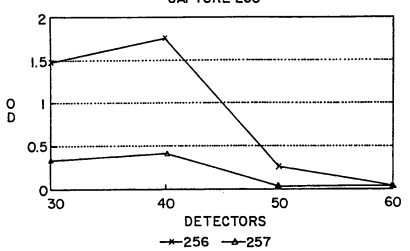
HPVI6 DETECTOR HYBRIDIZATION TEMPERATURE
CAPTURE 250



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Fig. 18

HPVI6 DETECTOR HYBRIDIZATION TEMPERATURE CAPTURE 250



PF	307	COCOL:	ENDPOI	/es in c nt				/IIX: OFF		WED. SEP 02 1992 4:53 PM CALIBRATION: ON			
1-6 96-91 1:200 OPTICAL DENSITY										6-12 1	37 . 91		
PEG	A	1 255-31 0.477	lH ₂		98		256	7 2.205	255		98		256
5% -T	В	0.036	0.033	0.032	0.032	0.038	0.038	0.036	0.039	0.035	0.036	0.037	0.037
1% BSA		0.418	0.500	0.240	0.349	0.155	0.128	2.709	2.003	0.839	0.551	2.051	1.932
·T	D	0.030	0.032	0.036	0.034	0.037	0.034	0.036	0.030	0.034	0.034	0.037	0.039
1%	BS	4						2.747					
							•	0.032					
0.19	6 P	VP 5						2.372					
-Τ	Н	0.036	0.035	0.034	0.030	0.030	0.034	0.032	0.031	0.032	0.032	0.033	0.044
ALL 250 AT 55°C 30' ALL IN 0.1% PVP, 5×5CC ALL IN 0.1% PVP, 5×33C DET AT RT IN GLYCEROL BUFFER DUPLICATE WELLS (-T = 96-91 OR 137-91 -T 3 5尺でより)													

	lOT	OCOL:	DETECT		OPTINI	ZATION AUTON					THU SEP 03 1992 11:33 AM				
WAV		ENGTH:		14.1		A01011	OII			CALIBRATION: 0					
1:200		—	256 -												
Final		1 0	2	3	4 5P/1BS	5 0	6 5%P		8 5P/1B	9 0	10 5%P	11 1%B	12 5P/1B		
96-9	iΑ	0.234	0.865	0.358	0.676	0.234	0.425	0.307	0.449	0.507	1.742	1.670	2.060		
٠ T	В	0.040	0.293	0.095	0.278	0.043	0.041	0.280	0.135	0.069	1.874	1.404	2.020		
	C	0.545	1.269	0.747	1.313	0.266	0.586	0.344	0.523	0.632	1.547	1.396	1.908		
.⊺	D	0.038	0.429	0.128	0.359	0.051	0.042	0.042	0.052	0.039	1.474	1.123	1.359		
	E	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000		
	F	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000		
	G	0.000	0.000	0.000	0.000	0.000	0.000	0.00	0.000	0.000	0.000	0.000	0.000		
	Н	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000		
STOP AT 2" CAPTURE 250, 55', 30', 0.1% PVP, 5×55 ^C PLATE 20' DET 0 = 30% GLYCEROL, 0.1% PVP, 1% BSA, 5×55 ^C 5%P = 5% PEG, 0.1%PVP, 1% BSA, 5×55 ^C 1%B = 1% BSA, 0.1% PVP, 5×55 ^C 5P/1B = 5% PEG, 1% BSA, 0.1% PVP, 5×55 ^C															

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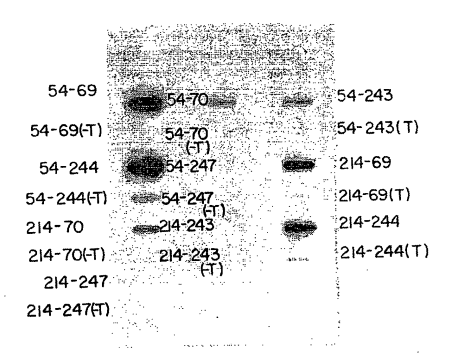


Fig. 22

RAW DATA

DES P	CRI RO	IPTION: TOCOL:	DATA01 HPV 18 ENDPO 450	CAPTU	RE AND		AUTON	NIX: ON				TUES. JAN 12 1993 7:46 PM Calibration: On		
			5	6	_		AL DENS	HY						
		1	2	3	4	267	6	7	8	9	10	11	12	
59	A	0.038	0.988	0.087	1.762	0.067	0.036	0.000	0.000	0.000	0.000	0.000	0.000	
260	В	0.033	1.129	0.033	2.621	0.037	0.031	0.000	0.000	0.000	0.000	0.000	0.000	
262	C	0.034	0.712	0.036	2.153	0.037	0.031	0.000	0.000	0.000	0.000	0.000	0.000	
268	D	0.037	0.919	0.037	2.311	0.038	0.037	0.000	0.000	0.000	0.000	0.000	0.000	
269	E	0.027	0.727	0.036	1.718	0.040	0.034	0.000	0.000	0.000	0.000	0.000	0.000	
270	F	0.026	0.237	0.038	0.662	0.040	0.030	0.000	0.000	0.000	0.000	0.000	0.000	
	G	0.034	0.037	0.036	0.040	0.034	0.033	0.000	0.000	0.000	0.000	0.000	0.000	
	Н	0.029	0.120	0.038	0.039	0.038	0.034	0.000	0.000	0.000	0.000	0.000	0.000	

Fig. 23

RAW DATA

DATA FILE: DESCRIPTION: PROTOCOL: MODE: WAVELENGTH:	HPV 16 .	AND 18	PLATE		AUTOM	IIX: ON	THU. JAN 14 1993 5:49 PM CALIBRATION: ON					
	CAPTURES						OPTICAL DENSITY 18 16					
1 BLANK a 0.038	2 267, 0.179	3 /250 0.208	-T ⁴ 0.035	0.035	6 267	-T '	8 2 0.365	9 .50 0.368	10 -T 0.046	98	12	
В	0.722	0.589	0.048	0.037	0.040	0.037	1.179	1.274	0.095	255		DE
С	0.454	0.408	0.036	0.041	0.049	0.040	0.778	0.754	0.059	256		DEFECTORS
D	2.367	2.429	0.035	2.619	2.626	0.038	0.039	0.043	0.040	260		T
E	2.607	2.593	0.035	2.724	2.695	0.039	0.527	0.524	0.038	98, 260		RS
F	2.842	2.742	0.047	2.729	2.773	0.040	1.427	1.537	0.174	255, 260		
G	2.781	2.799	0.043	2.894	2.804	0.097	1.034	1.054	0.125	256, 260		
Н	0.041	0.042	0.038	0.042	0.042	0.044	0.039	0.056	0.043			

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Fig. 24

HPV EPA
CONCENTRATION VS SIGNAL

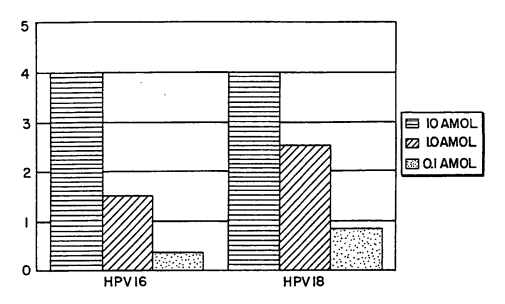
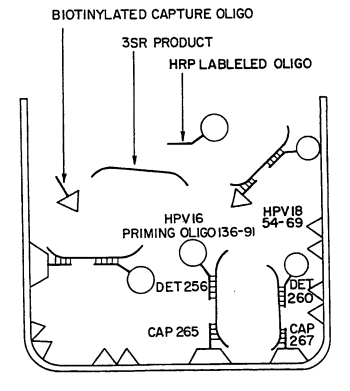
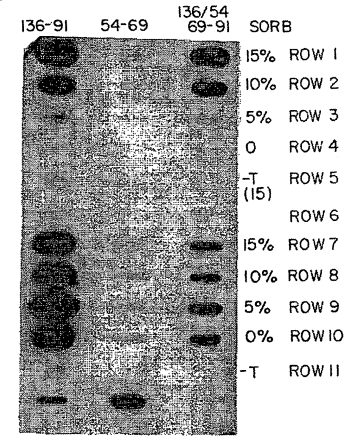


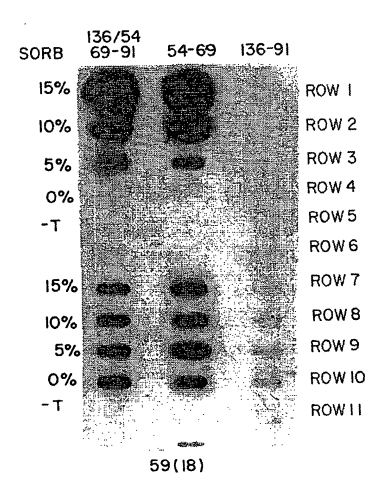
Fig. 25



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STREPTAVIDIN





SUBSTITUTE SHEET (RULE 26)